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# **Tritium Inventory in ARIES-AT**

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### **ABSTRACT**

This report documents an investigation into the tritium inventory expected in the ARIES-AT fusion reactor. ARIES-AT features silicon carbide fibers in a silicon carbide matrix as its primary construction. It uses the same fusion power core as the previous ARIES-RS. Based on experimental results of several researchers, consideration was given to swelling, sputtering, film coatings, erosion, and implantation. Estimates were made of tritium inventory using the TMAP4 code. About 700 g of tritium may be expected in the machine, two thirds of which would reside in the first wall. Under assumed accident conditions that involve firs wall temperatures up to  $1000^{\circ}\text{C}$ , evolution of retained tritium may be expected to vary from 0.8 to nearly 40 percent depending on the temperature of the first wall.

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## **Tritium Inventory in ARIES-AT**

#### **OBJECTIVE**

The objective of this study is to estimate tritium inventory in ARIES-AT and to evaluate safety consequences associated with that inventory. The approach makes use of available design information, estimates of surface particle fluxes and energies, and the TMAP4 code to assess the buildup of tritium in the structure and, to the extent possible, in the cooling system. Then some accident scenarios will be investigated to evaluate tendencies for tritium release.

#### MACHINE DESCRIPTION

As presently configured, ARIES-AT makes use of essentially the same power core configuration as the previous ARIES-RS, but the structure is different. Figure 1 shows the general layout of the machine. The plasma has a double-null divertor configuration and a net plasma fusion power of 2170 MW.

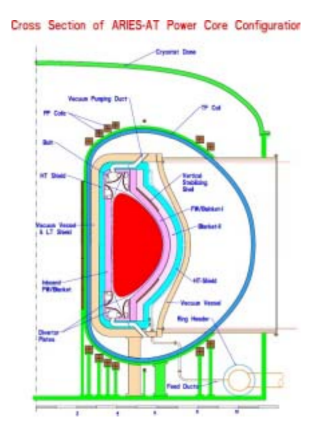
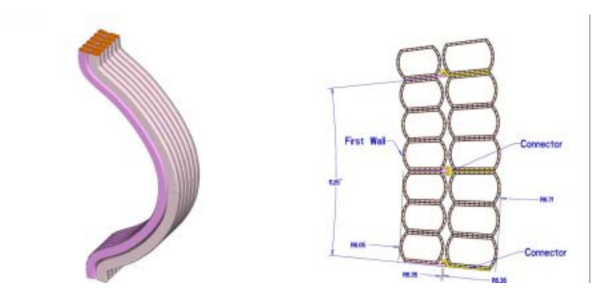


Figure 1. General layout of ARIES-AT

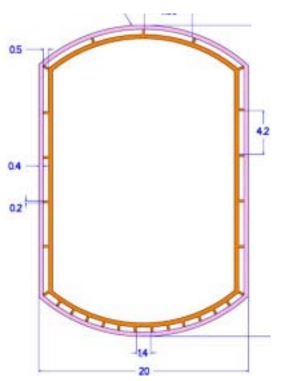
Key design parameters significant to tritium inventory are the construction of the first wall-blanket (FW/B) and divertor sections, the coolant flows and the plasma particle fluxes to the adjacent structures. Figure 2 shows the general structure of the segments making up the first wall-blanket, and Figures 3 and 4 show additional details. Figure 5 shows some detail for the divertor region.

The plasma-facing structure of ARIES-AT is of  $SiC_f/SiC$  (silicon carbide fibers woven and embedded in a silicon carbide matrix). The structures are rather thin but stiff, essentially concentric shells connected by thin webs. The divertor plates are of tungsten.

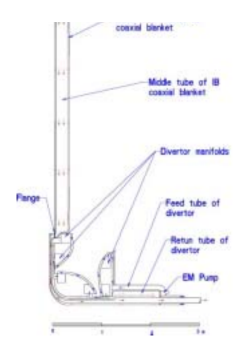


**Figure 2.** ARIES-AT outboard first wall and blanket.

Figure 3. Outboard first wall/blanket cross section.



**Figure 4.** ARIES-AT first wall configuration details.



**Figure 5.** ARIES-AT divertor coolant flow paths.

#### TRITIUM INVENTORY

#### Radiation Effects

The anticipated lifetime of the FW/Blanket structure is 3 years. With a nominal 14-MeV neutron power of 1390 MW over 430 m<sup>2</sup> of surface area, the average neutron flux rate would be  $1.4 \times 10^{18}$  n/m<sup>2</sup> s. If there were 300 effective full power days per year, and there are 1.16 dpa/ $10^{25}$  n/m<sup>2</sup>, then FW structures may be expected to have about 12.5 dpa at the end of their lifetime. That may generate significant changes in the material. Here we consider what some of those changes may be.

ARIES-AT will operate at elevated FW temperatures. The ARIES-AT requirement is that the SiC operate at less than 1000°C. Present estimates are that the FW will operate with a maximum surface heat flux to the wall of 0.7 MW/m² with an average surface heat flux of 0.5 MW/m² [¹]. Thermal conductivity for the SiC<sub>f</sub>/SiC material varies with the method of production, density, degree of damage from irradiation, and other factors. It could be as high as 30 W/m K [²] at the working temperature, but a more realistic value may be 20 W/m K [1]. The design-team calculated temperature distribution in the FW structure is peaked at about 1000°C near the midplane, falling off to about 700°C at the bottom end and to about 850°C at the top.

A further consideration is the potential for annealing of defects and recrystallization of the SiC at elevated temperatures. Kawatsura et al. <sup>3</sup> have found that the critical dose of implanted ions for amorphization of the SiC crystal depends on the mass of the implanting ion and on temperature. They found that at room temperature, the critical dose best-fit line varies according to

$$\Gamma_{crit} \left( \frac{atom}{m^2} \right) = 2.2 \times 10^{20} \ Z^{-1.3} \tag{1}$$

meaning that for H (Z = 1), the critical dose is  $2.2 \times 10^{20}$  atom/m<sup>2</sup>. Kawatsura et al. cite data from Spitznagel et al. [<sup>4</sup>] giving the critical flux for H into SiC as  $9 \times 10^{20}$  atom/m<sup>2</sup>. Presumably, the Spitznagel data are for ambient-temperature implantation.

In annealing investigations, Kawatsura et al.  $^3$  observed a 25-nm  $SiO_2$  layer on an  $\alpha$ -SiC crystal surface after annealing for 2 h in argon at 1200°C. Such films may retard sublimation of SiC at elevated temperatures, and they may themselves sublime.  $^5$  Oxygen incorporated in the fibers may also come out as CO.

No change in the crystal structure (amorphization) was seen after a 2-h anneal at 1000°C. After annealing at 1500°C, however, Kawatsura et al. observed a defect layer with a very high concentration of the implanted Ni atoms at the 540-nm implantation depth. For ARIES-AT, the implantation depth is expected to be in the range of a few tens of angstroms. It is not clear to what extent the damage region would develop for such shallow implantation depths.

Neutron irradiation produces linear swelling in SiC. Scholz<sup>7</sup> quoting Price<sup>8</sup> indicates that swelling saturates at about 0.5 dpa, and the amount of swelling at saturation varies with temperature according to

$$\frac{\Delta L}{L} = 0.011 - 1.1 \times 10^{-5} \, T(^{\circ}C) \tag{2}$$

The data indicate that for an operating temperature of 1,000°C, one may expect linear swelling to saturate at about 0.1 percent, though for regions where the temperature is lower, the swelling may reach 1 percent or so.

Swelling may have an effect on tritium retention, particularly if the increased volume is partially filled with tritium atoms or molecules. Saito et al.  $^9$  found that implantation of He at 3 MeV to about 10,000 appm produced bubbles that were observable after annealing at 1,673 K, but not before annealing. These bubbles appeared in the SiC matrix only and not in the SiC fibers. Scholz et al.  $^{10}$  injected He to 2,500 appm in SiC composites at 1175 K. They observed swelling of the matrix but contraction of the fibers. They concluded that the presence of the He may suppress annealing of structural damage at that temperature. Similar results were seen by Frias Rebelo et al.  $^{11}$  where the effect was seen after both neutron and helium ion exposure. Bacon et al.  $^{12}$  reported gas-driven flaking when SiC at 700°C was bombarded with 40-keV helium ions to a dose of  $5 \times 10^{21}$  ion/m $^2$ . Flake thickness corresponded to the implantation depth, and its existence was thought to correspond to the absence of amorphous structure at elevated temperatures.

Another issue of concern is the potential for composition change at the surface due to sputtering actions. Carbon and silicon atoms are expected to be sputtered individually. In the presence of hydrogen, carbon sputtered from a carbon substrate is known to redeposit as amorphous hydrocarbon  $CH_x$ , where x is in the range of 0.1–0.4, the value depending on temperature, hydrogen partial pressure, and possibly other factors. Silicon is also believed to form hydrogenic films. It is not known to what extent these processes would be applicable in ARIES-AT.<sup>13</sup> Data that may be relevant are from Xu et al.<sup>14</sup> who observed that vapor deposition of SiC from methyltrichlorosilane in a hydrogen gas carrier at temperatures ranging from 1000 to 1300 °C resulted in the formation of pure, mostly  $\beta$  phase SiC crystallites. Neither free silicon nor free carbon was found. The SiC was poorly crystallized and very fine (~10 nm). As the deposition temperatures decreased, the crystallite size became smaller. They also observed the formation of a thin (24-nm) SiO<sub>2</sub> film from system residual oxygen. That film was readily sputtered away under the Auger analysis beam in about 10 minutes. Lower surface temperatures may favor amorphous hydrogenic film formation whereas at higher temperatures it may not form.

Mohri et al. <sup>15</sup> measured sputtering rates of SiC bombarded with  $H^+$ ,  $D^+$ , and  $Ar^+$  ions at energies of 5-13 keV and various incidence angles. They found erosion yields for  $H^+$  incident at an angle of 45° peaked at about 8 keV with a value of about 0.03 atoms/ion. Angular dependence showed the greatest yields at angles from 50-80°. They also saw evidence for the formation of hydrocarbons and silicon hydrides during sputtering with  $H^+$ . Sputtering yields for  $D^+$  were approximately a factor of 2 higher than for  $H^+$ , as may be expected. They observed that

sputter yields are greater for fully dense material, such as they used, than for porous material. This is apparently due to the capacity of the porosity to port off implanted gas atoms that in fully dense material cause blistering and flaking.

These considerations lead to the following summary of possibilities for modeling, regarding the state of the  $SiC_f/SiC$  in the ARIES-AT environment.

- There may be a thin SiO<sub>2</sub> film on the SiC surface, but more likely, it will be sputtered clean. Oxide films have been observed on many materials in similar situations. There may well be a competition between film formation and film erosion due to the sputtering effect of the plasma that will depend on temperature and on the oxygen impurity content of the plasma. If such films form and inhibit return of implanted hydrogen isotope atoms to the plasma, continued implantation will lead to exceptionally high concentrations of implanted atoms at the implantation depth that will probably induce flaking.
- It is also possible that there will be thin films of amorphous C:H and Si:H. Hydrogenic films are expected to be more permeable to hydrogen isotopes than SiC will be. Therefore, if any such films grow to thicknesses exceeding the implantation depth, implantation burden to the SiC substrate would stop.
- Neutron irradiation may generate trapping sites in the SiC. Swelling should saturate within about 4 percent of the FW lifetime, so it may be possible to simply model traps as constant at about 1,000–10,000 appm. Trap strength is not known, but it may be about 2 eV, the molecular formation free energy per atom. Traps would be more or less uniformly distributed throughout the thickness of the wall.
- The tendency of the matrix SiC to swell while the fibers contract under irradiation has the potential for partial debonding of the matrix from the fibers. If this happens, there may be microchannels for release of tritium and other gases along the interface between the matrix and the fibers. If that is true, then the mean diffusion distance for release of implanted tritium would be only about the fiber diameter or about 10 µm. On the other hand, tritium coming from an effective partial pressure at the surface of the SiC could migrate through the structure much faster. The effects of these channels could be modeled by using a higher diffusivity.

## Plasma Particle Loading in ARIES-AT

There have been few studies of plasma edge conditions and particle fluxes to surfaces for the ARIES-AT design. Obviously, the device is a tokamak, and the plasma should exhibit conditions similar to those of any other large power tokamak concept studied by the community thus far. However, little work has been done yet specifically to address plasma properties, especially edge plasma issues, for the ARIES-AT. One of the greatest areas of uncertainty is the degree of detachment of the plasma from the divertor. That will have a profound effect on the particle flux density and particle energies impacting the divertor plate.

Plasma conditions are believed to be comparable to those postulated for the International Thermonuclear Experimental Reactor (ITER)<sup>13</sup>. Those conditions are summarized in Table 1. The various areas referred to in Table 1 are shown schematically in Figure 6.

**Table 1**. Plasma particle loads estimated for ITER and thought to be representative of those that may be seen in ARIES-AT. <sup>16</sup>

Region	Particle Flux Density (10 <sup>20</sup> DT/m <sup>2</sup> s)	Particle Characteristic Energy (eV)	Fast Neutron Flux Density (10 <sup>18</sup> n/m <sup>2</sup> s)
First Wall	0.1–1	100–500	1.9–2.3
Limiter	10–100	100-500	2.3
Upper Baffle	0.1–1	100–500	2
Lower Baffle	1–10	100	1.1
Lower Target	<10,000	<5	0.4-0.6
Divertor Sidewall	1–10	5–100	0.6–1.1
Dome	1-10	5–100	0.6–1.1

In addition to particle streaming into the plasma-facing surfaces, mostly as charge-exchange neutral atoms, there will be some tendency of neutral gas molecules to circulate around the structures and become incorporated into surfaces by codeposition or simple plate out. Such was seen in both TFTR and JET where inventories of tritium were high in the side surfaces of tiles and in carbonaceous flakes deposited on and near vacuum pumping port louvers. <sup>17,18</sup>

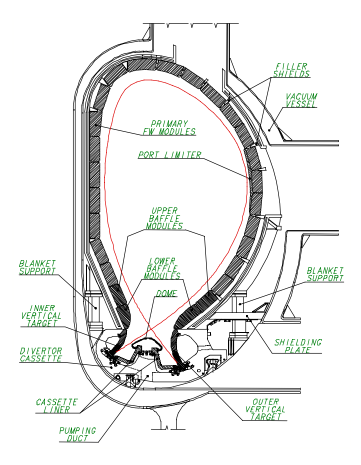
TRIM calculations suggest average implantation depths for normal incidence of D/T in SiC varying from 35 angstroms at 100 eV to 125 angstroms at 500 eV ion energy. Straggle varies correspondingly from 20 to 60 angstroms. Those numbers don't change much if there is a 25-angstrom surface layer of SiO<sub>2</sub>.

## **TMAP Modeling**

The recommended<sup>19</sup> value for solubility of hydrogen isotopes in SiC is

$$S\left(\frac{\text{atom}}{\text{m}^{3}\text{Pa}^{1/2}}\right) = 2.58 \times 10^{19} \exp\left(\frac{0.61 \text{ eV}}{\text{kT}}\right)$$
(3)

while hydrogen diffusivity is



**Figure 6**. Cross section of ITER plasma chamber showing various regions referred to in Table 1 for plasma particle fluxes. <sup>16</sup>

$$D\left(\frac{m^2}{s}\right) = 9.8 \times 10^{-8} \exp\left(-\frac{1.89 \text{ eV}}{kT}\right)$$

We can explore several bounding conditions on implantation uptake for SiC in the first wall. One is the assumption that there is enough surface activity of hydrogen isotopes and other species, or sufficiently high porosity in the surface, that there is a zero surface concentration boundary with implantation at the nominal depth and scatter, and assume that there are no traps. A second option would assume traps at 10,000 ppm (0.01 atom fraction) concentration and 2.1 eV trapping energy. This is close enough to the diffusion activation energy that it will not have much effect. A boundary where there is very little reemission (non-flow boundary) is unlikely because the extremely high loading at the implantation depth would promote blisters or flaking. Another option might be a Sieverts' law boundary condition, but estimates of gas pressures near the surfaces would be needed to render that assumption viable. Advice<sup>13</sup> is that there have not yet been enough edge plasma studies to justify guesses on those pressures.

The TMAP4 code was used to get some quantitative estimates of tritium inventory buildup. For simplicity, the computations were conducted for deuterium alone rather than a mix of

deuterium and tritium, but it is not significantly different than if a D/T mix were used. For this segment of the analysis, the plasma facing material is assumed to be purely SiC and to consist of a simple slab 4 mm thick. Appendix A contains several TMAP input files and selected results from that modeling.

The first or reference case used an implantation flux of  $10^{21}$  D/m²-s with implantation distributed approximately as calculated by TRIM. Both upstream and downstream faces were assumed to be at the "c = 0" condition for diffusive boundary conditions. The inventory after three years was only about 7 percent (1.75 g/m²) of its saturation value (24.7 g/m²). Increasing the implantation flux by a factor of ten increased the inventory by the same factor of ten.

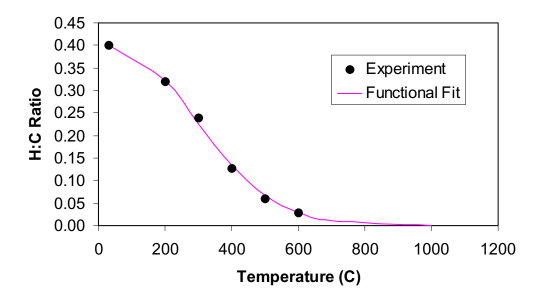
The second case modeled was similar to the first, but it included a temperature gradient of 200 K across the 4-mm thickness of the sample, beginning at the same 1,000°C at the implantation face as in the previous case. This gradient is probably an extreme, but it was selected to evaluate the effect. Also included in another simulation for comparison were traps at 2.1 eV trap energy and 0.01 atom fraction concentration. The temperature gradient added almost a factor of 10 to the inventory by inhibiting diffusion through the bulk of the wall and forcing higher concentrations near the implantation zone. These reached to about half of the SiC lattice atom density in the implantation region. The traps added 1 percent to the total inventory, as may have been be predicted *apriori*. Again, further details are in Appendix A.

Another inventory of implanted tritium will be in the divertor plates. Because they are of tungsten, as were the divertor plates in the ITER concept studied previously, <sup>20,21</sup> with effectively the same loading and operating conditions (at least no significant differences have been yet identified), we may be justified in simply bringing those results forward to this study. The estimated specific inventory in the 10-mm thick tungsten sidewall divertor plates in ITER was 0.4 g/m<sup>2</sup>. For a divertor area of 67.7 m<sup>2</sup>, that translates to a 27 g inventory in ARIES-AT. Even though particle flux densities are much higher to these components than to the first wall, the saturation effect in tungsten <sup>22</sup> and high recycling means that these structures will not have so much tritium burden as will the SiC structures. This inventory was calculated to saturate in less than 0.1 years of operation.

## **Codeposition Estimates**

Codeposition estimates are essentially wholly dependent on the rate of sputtering of carbon and silicon from the SiC surfaces. The carbon atoms will be reionized and redeposit in a short distance, a very few cm, from the surface. The ionization mean free path for silicon atoms is much longer, leading to the suspicion that amorphous carbon and silicon films may form under tokamak conditions. Experience has shown that these amorphous film molecules tend to migrate to the coolest surfaces they have access to, which often is on the sides of plasma-facing structures or vacuum pumping ducts.

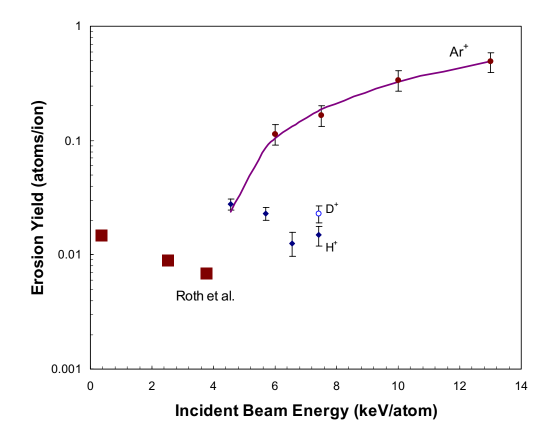
The hydrogen content of codeposited hydrocarbon films depends on the temperature at which they are formed. Results from Doyle<sup>23</sup> on the hydrogen content of such films as a function of temperature are shown in Figure 7 together with a functional fit to those data. It may be seen there that if the operating temperature is up around 1,000°C, the H:C ratio will be low.



**Figure 7.** Experimental data from Doyle et al.<sup>23</sup> for the H:C ratio as a function of formation temperature, and a functional fit to those data.

The functional fit suggests it will be of order 2 x 10<sup>-4</sup>. At 800°C it is about 3 x 10<sup>-3</sup>. As noted earlier, the design-team calculated temperature distribution in the FW structure is peaked at about 1000°C near the mid-plane, falling off to about 700°C at the bottom end and about 850°C at the top. The non-linearities shown in Figure 7 imply that the greatest tritium inventory in codeposited films will be in those areas that are the coolest. For an order of magnitude estimate, suppose that the mean effective temperature of this coolest area is 700°C and that it represents 20 percent of the first wall area or about 60 m<sup>2</sup>. The functional fit in Figure 7 suggest a H:C ratio of 0.01 at 700°C. If tritium constitutes half the hydrogen, then T:C is about 0.005 at that temperature.

Figure 8 shows sputter yield data from Mohri et al<sup>15</sup> for H<sup>+</sup>, D<sup>+</sup>, and Ar<sup>+</sup> bombarding SiC. These are shown in terms of energy per H or Ar nucleus based on reported fractions of H, H<sub>2</sub><sup>+</sup>, and H<sub>3</sub><sup>+</sup> and the stated beam energy. For comparison, similar data by Roth et al.<sup>24</sup> are shown. The right-most data point from Roth et al. was for less than fully dense material, while the rest of the data were for fully dense SiC. Additionally, the experiments of Roth et al. were at normal incidence whereas Mohri et al. conducted their experiments at 45° incidence angle where yields should be higher. The figure suggests that sputter yield of SiC will be about 0.01 regardless of isotope or energy. Less than fully dense material appears to experience less sputtering because of the propensity for the voids to port away hydrogen that in fully dense material would cause micro-blistering and contribute to the erosion. There was no apparent chemical erosion of the SiC due to the H or D ions, whereas in graphite sputtering, chemical erosion dominates in some temperature regimes.



**Figure 8.** Erosion yields as a function of incident beam energy at 45° incidence from Mohri et al. <sup>15</sup> Also shown are normal-incidence data for H<sup>+</sup> bombardment of SiC from Roth et al. <sup>24</sup> The line through the Ar<sup>+</sup> data is a least-squares linear fit.

If we multiply the  $330\text{-m}^2$  first wall area by an assumed particle flux of  $10^{20}$  D/T/m<sup>2</sup>s (Table 1) for three years, we estimate the amount of SiC sputtered to be  $5 \times 10^4$  moles. With a H:C ratio of 0.01 (H:Si may be the same or slightly lower based on formation enthalpies) and assumed full deposition of the material in the vacuum vessel or in pumping ducts, the tritium inventory would be 50 moles, 150 g, or 1.5 MCi, assuming all of the sputtered carbon and silicon combine in the ratio stated and that half of the codeposited species would be D and half T. This is less than but comparable, considering the gross approximations made here, with the 580 g estimated by TMAP calculations in the next section to be resident in the first wall. The 150 g is about 0.2 percent of the tritium striking the first wall.

## **Order of Magnitude Inventory**

An order of magnitude estimate of tritium inventory in the ARIES-AT may be drawn from the above considerations to be as shown in Table 2. It will be seen there that the preponderance of the nominal 760-g in-vessel inventory is in the SiC first wall structure, resident in the SiC itself. That grows nearly linearly in time and is limited by the low diffusivity of hydrogen

species in the SiC. The next largest inventory is in the codeposited species that are assumed to plate out on the colder surfaces of the vacuum vessel or surrounding duct work. This estimate is highly uncertain because of the uncertainty in the tritium fraction in redeposited silicon and carbon, but also because of the uncertainty in the sputter yields for the plasma facing structures. Tritium in the divertor is least significant, but probably the best known.

**Table 2**. Order-of-magnitude estimate of tritium inventory in ARIES-AT after 3 years of operation. Figures could be low or high by a factor of 2.

Structure	Inventory (g)	
First Wall	534	
Divertor	27	
Codeposition Regions	150	
Total	711	

It should be emphasized that these are single point values without the 2- or 3-dimensional nature of the various processes or states that influence tritium inventories taken into account.

#### **ACCIDENT CONSEQUENCES**

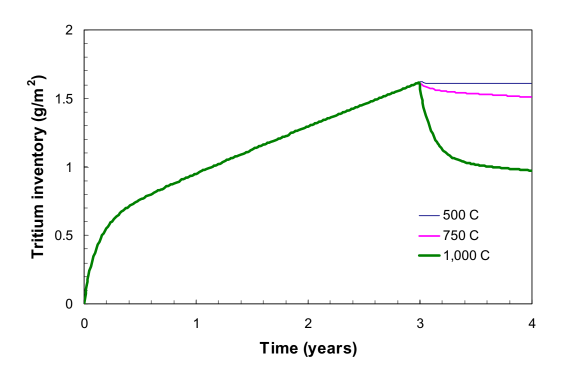
Suppose there were a loss-of-vacuum accident in which the vacuum chamber were to be penetrated with the associated development of an opening to the surroundings. The greatest immediate concern would be the levitation of the codeposited material and dust in the vessel and its potential for being carried to the surroundings by connective flows of air. The tritium inventory fraction so released would depend on the location of the break relative to the location where the codeposition took place and on the chemical state of the codeposited material. In some machines, the codeposited material became flaky and had a rather large dust component associated with it. In other machines, or in other parts of the same machines, there are areas where the codeposited film is tightly bonded to the first wall as a carbonaceous or polymer-like film. The former is subject to levitation, while the latter generally is not. Exposure to air, particularly if the coated walls are at elevated temperatures and/or the air is humid, can degrade the integrity of those coatings and allow the films to loosen and be eroded. That may be a concern over lengthy outages due to accidents.

Tritium in the tungsten divertor plates will come out at a high rate, even if the plates are at ambient temperature. Hence, any time the machine is non-operational, one may expect to see tritium coming from the divertor plates. This would constitute a continuing source if the machine were left open for any significant time with much of the inventory in the tungsten

coming out in days to weeks, depending on the temperature. If the machine were not open but unpumped, the inventory of gaseous tritium in the vacuum chamber would climb.

It seems reasonable that with the bulk of the structure in the vacuum chamber being SiC, there would not be a large amount of structural afterheat from radioactive decay, except possibly for the tungsten divertor plates. Hence, under accident conditions, it is likely that the temperatures of the SiC structures would not exceed their normal operating temperatures. The time it took to build the inventory in the walls would then be required to have the tritium escape, even if there were a readily absorbing medium at the SiC surface, such as moist air.

To evaluate that, we consider several accident scenarios using the TMAP4 code. First, the inventory is allowed to build for three years of operation under the worst-case conditions for the first wall, namely that there is effectively no reemission. Then it is assumed there is an accident, and the wall temperature is assumed to remain at one of several arbitrary temperatures for one year under the new worst-case scenario, that of essentially zero concentration at the first wall surface. Thus, the release is dominated by diffusivity in the SiC. Following the accident, the SiC structure is assumed to be held at 500, 750, and 1,000°C. Release fractions after one year are 0.8, 6.6, and 39.7 percent, respectively. Results are shown in Figure 9



**Figure 9**. Release of tritium following three years of operation at  $10^{21}$  DT/m<sup>2</sup>.s and assuming SiC structure is held at 500, 750, and 1,000°C following an accident. Release fractions after one year are 0.8, 6.6, and 39.7 percent, respectively.

If there were to be a major thermal excursion that could raise the SiC temperature to 1,400 or 1,600°C, there would probably be a significant change in the microstructure of the SiC. Bubbles would grow as swelling took place. There would be some connection of the bubbles with the surface through porosity that would develop, and the majority of the tritium inventory in the SiC would emerge in a short time.

Specifics of accidental releases must await better definition of conditions attendant to such accidents.

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### APPENDIX A: TMAP INPUT FILES AND RESULTS

### **Reference Case**

Input file and inventory buildup history for zero concentration boundary conditions, where SiC is implanted at 100 eV, normal incidence at a flux density of  $1.0 \times 10^{21} D/m^2$ -s. Results are generally representative of a D/T mixture. After nominally 149 weeks, the inventory was  $7 \times 10^{23} D/m^2$ . If half of that inventory were tritium, that would correspond to  $1.75 g/m^2$  of surface area. Increasing the implantation flux by a factor of ten increases the inventory by that same factor of ten.

```
title input
ARIES AT First Wall, 5-mm SiC, Surf Temps 1000 C, d only - implanted
D+T flux=1.0e21/m2s, Causey properties, C=0 BCs
end of title input
main input
dspcnme=d, end
espcnme=d2, end
segnds=7,12,end
nbrencl=2,end
linksegs=1,2,end
end of main input
$ -----
enclosure input
$ -----
start bdry,1,end $upstream side
$ Enclosure 1 is the plasma side
etemp=1273.,end
esppres=d2, const, 0.0, end
start bdry,1,end
                       $upstream side
$ Enclosure 2 is the coolant side
etemp=1000.,end
esppres=d2, const, 0.0, end
end of enclosure input
$ -----
thermal input
$ -----
start thermseg, end
$ SiC implantation zone
delx=0.,5*2.e-9,0.,end
                         $ Damage/implant thickness 100 Angstroms
dtemp=7*1273.,end
                                   $ Constant temperatures=(K)
start thermseg, end
$ Bulk SiC
delx=0.,1.e-9,1.e-8,1.e-7,1.e-6,1.e-5,1.e-4,4*1.e-3,0.,end $ 4 mm thick
dtemp=12*1273.,e
                                   $ Initial temperatures=(K)
```

```
end of thermal input
diffusion input
start diffseq,end
$ SiC implantation zone, 100 Angstroms
dconc=d,7*0.,end
                              $ Starting mobile concentration
qstrdr=d,const,0.,end
                              $ Q*/R for Soret effect
{\tt spcsrc=d,tabl,1,srcpf,0.,.263,.351,.263,.096,.026,0.0,end}
dcoef=d,equ,1,end
                             $ diffusion coeff (m2/s)
difbcl=sconc,encl,1,d,d2,nsurfs,2,conc,const,0.0,end
difbcr=link,d,solcon,equ,2,end
surfa=1.0, end
$
start difseg, end
$ Bulk SiC
dconc=d,12*0.,end
                                  $ Starting mobile concentration
                                  $ Q*/R for Soret effect
qstrdr=d,const,0.,end
spcsrc=d,const,0.,srcpf,12*0.,end
dcoef=d,equ,1,end
                                  $ diffusion coeff (m2/s)
difbcl=link,d,solcon,equ,2,end
difbcr=sconc,encl,1,d,d2,nsurfs,2,conc,const,0.0,end
surfa=1.0,end
end of diffusion input
equation input
$(1) Diffusivity of hydrogen in SiC (Causey)
y=9.8e-8*exp(-1.98/8.625e-5/temp), end
$(2) Causey's solubility
y=2.58e19*exp(0.61/8.625e-5/temp),end
end of equation input
table input
$ Implantation flux
0.0,1.e21,1.e8,1.e21,end
end of table input
control input
time=0.,end
tstep=100.,end
                                      $ time step = 100 sec
                                      $ end at time = about 3 years
timend=9.e7,end
                                      $ print every week
nprint=6048, end
itermx=900,end
delcmx=1.e-5,end
end of control input
plot input
nplot=6048,end
                      $ makes plotfile entry every week
plotseg=1,2,end
                      $ segments for which plot info is needed
plotencl=1,2,end
                      $ enclosures for which plot info is needed
dname=d,end
                      $ diffusing species for which plot info is needed
ename=d2,end
                      $ enclosure species for which plot info is needed
dplot=moblinv,sflux,end
eplot=diff,end
                      $ flow of molecules into enclosure
end of plot input
```

\$
end of data

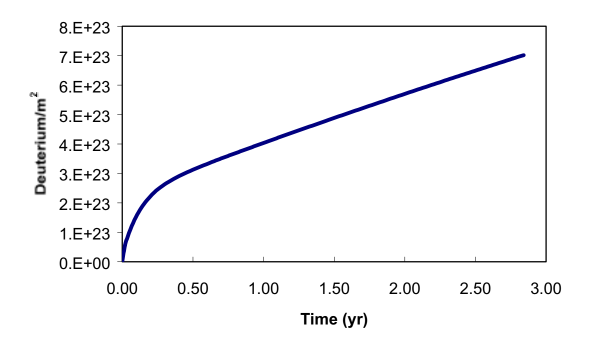


Figure A-1. Calculated inventory buildup history for deuterium implanted at 100 eV with zero concentration boundary conditions. Assymptotic inventory limit is  $4.96 \times 10^{24} \, (\text{D/m}^2)$ 

## With Temperature Gradient and Trapping

```
title input
ARIES AT First Wall, 5-mm SiC, Surf Temps 1000 C, d only - implanted
D+T flux=1.0e21/m2s, Causey properties, nonflow left BC
end of title input
main input
dspcnme=d, end
espcnme=d2, end
segnds=7,12,end
nbrencl=2,end
linksegs=1,2,end
end of main input
enclosure input
start bdry,1,end
                           $upstream side
$ Enclosure 1 is the plasma side
etemp=1273.,end
esppres=d2, const, 0.0, end
start bdry, 2, end
                           $downstream side
$ Enclosure 2 is the coolant side
etemp=1000.,end
```

```
esppres=d2,const,0.0,end
end of enclosure input
$ -----
thermal input
$ -----
start thermseq, end
$ SiC implantation zone [thermseg 1]
delx=0.,5*2.e-9,0.,end
                       $ Damage/implant thickness 100 Angstroms
dtemp=7*1273.,end
                                      $ Constant temperatures=(K)
tcon=const,20.0,end
rhocp=const, 4.7e7, end
hsrc=const,0.0,srcpf,7*0.0,end
htrbcr=link,end
htrbcl=stemp,const,1273.,end
hgap=const,1.e6,end
$
start thermseg, end
$ Bulk SiC [thermseg 2]
delx=0.,1.e-9,1.e-8,1.e-7,1.e-6,1.e-5,1.e-4,4*1.e-3,0.,end $ 4 mm thick
dtemp=12*1273.,end
                                         $ Initial temperatures=(K)
tcon=const, 20.0, end
rhocp=const, 4.7e7, end
hsrc=const,0.0,srcpf,12*0.0,end
htrbcl=link,end
htrbcr=stemp,const,1073.,end
end of thermal input
diffusion input
start diffseg, end
$ SiC implantation zone, 100 Angstroms [diffseg 1]
dconc=d,7*0.,end
                            $ Starting mobile concentration
qstrdr=d,const,0.,end
                             $ Q*/R for Soret effect
spcsrc=d,tabl,1,srcpf,0.,.263,.351,.263,.096,.026,0.0,end
dcoef=d,equ,1,end
                             $ diffusion coeff (m2/s)
ctrap=d,7*0.0,end
trapping=cetrpi,0.01,nbrden,4.833e28,d,alpht,equ,3,alphr,equ,4,end
difbcl=sconc,encl,1,d,d2,nsurfs,2,conc,const,0.0,end
difbcr=link,d,solcon,equ,2,end
surfa=1.0, end
$
start difseq, end
$ Bulk SiC [diffseg 2]
                                 $ Starting mobile concentration
dconc=d,12*0.,end
qstrdr=d,const,0.,end
                                 $ Q*/R for Soret effect
spcsrc=d,const,0.,srcpf,12*0.,end
dcoef=d,equ,1,end
                                 $ diffusion coeff (m2/s)
ctrap=d,12*0.0,end
trapping=cetrpi,0.01,nbrden,4.833e28,d,alpht,equ,3,alphr,equ,4,end
difbcl=link,d,solcon,equ,2,end
difbcr=sconc,encl,2,d,d2,nsurfs,2,conc,const,0.0,end
surfa=1.0,end
end of diffusion input
```

```
equation input
$(1) Diffusivity of hydrogen in SiC (Causey)
y=9.8e-8*exp(-1.98/8.625e-5/temp), end
$(2) Causey's solubility
y=2.58e19*exp(0.61/8.625e-5/temp),end
$(3) Trapping rate equation
y=1.3e12*exp(-1.98/8.625e-5/temp), end
$(4) Release rate equation
y=1.e13*exp(-2.13/8.625e-5/temp), end
end of equation input
table input
$ Implantation flux
0.0,1.e22,1.e8,1.e22,end
end of table input
control input
time=0.,end
tstep=100.,end
                                      $ time step = 100 sec
timend=9.e7,end
                                      $ end at time = 3 years
nprint=6048,end
                                      $ print every week
itermx=900,end
delcmx=1.e-5,end
end of control input
plot input
nplot=6048,end
                      $ makes plotfile entry every week
                      $ segments for which plot info is needed
plotseg=1,2,end
plotencl=1,2,end
                      $ enclosures for which plot info is needed
                      $ diffusing species for which plot info is needed
dname=d,end
ename=d2,end
                      $ enclosure species for which plot info is needed
dplot=moblinv,sflux,end
                      $ flow of molecules into enclosure
eplot=diff,end
end of plot input
end of data
```

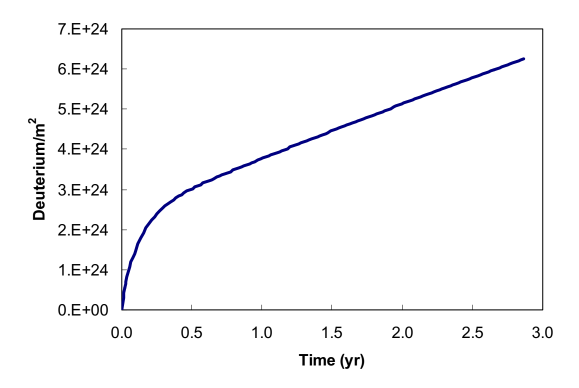


Figure A-2. Calculated inventory buildup history for deuterium implanted at 100 eV with zero concentration boundary conditions, but modified with thermal gradient and trapping. Traps with 2.1-eV trap energy and 0.01 atom fraction added 1 percent to the inventory.